QUANTITATIVE PREDICTION METHOD

The present invention concerns methods and systems for analysis of drug resistance in HIV-1. More specifically, the invention provides methods for predicting drug resistance by correlating genotypic information with phenotypic profiles. The methods allow the identification of primary and secondary resistance-associated mutations for new and existing drugs and for calculating the contribution of mutations and combinations of mutations to resistance and hyper-susceptibility. The invention allows the design, optimization and assessment of the efficiency of a therapeutic regimen based upon the genotype of the disease affecting a patient.

This application claims priority benefit of EP patent application nr. 03101687.6, and of U.S. Provisional Application No. 60/478,780 filed on June 16, 2003, the contents of which are expressly incorporated by reference herein. All other publications, patents and patent applications cited herein are incorporated in full by reference.

15 BACKGROUND

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Techniques to determine the resistance of HIV-1 to a therapeutic agent are becoming increasingly important. Many patients experience treatment failure or reduced efficacy over time. This is generally due to the virus mutating and/or developing a resistance to the treatment. As used herein, "HIV" is the human immunodeficiency virus, which is a retrovirus.

The various different anti-HTV-1 agents that have been developed over the years were initially administered to patients alone, as monotherapy. Though a temporary antiviral effect was observed, all the compounds lost their effectiveness over time. Research has now demonstrated that one of the main reasons behind treatment failure for all the antiviral drugs is the development of resistance of the virus to the drug (see, for example, Larder et al., 1989, Science, 246, 1155-8). This is largely due to the ability of HTV continuously to generate a number of genetic variants in a replicating viral population. These genetic changes generally alter the configuration of the HTV reverse transcriptase (RT) and protease (PR) molecules in such a way that they are no longer susceptible to inhibition by compounds developed to target them. If antiretroviral therapy is ongoing and if viral replication is not completely suppressed, the selection of genetic variants is inevitable and the viral population becomes resistant to the drug. Since then, dual combination therapy, using drugs that target both HTV reverse transcriptase (RT) and protease (PR) molecules, has provided increased control of viral replication, and thus provided extended clinical benefit to patients. In recent years,

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however, it has become clear that even patients being treated with triple therapy including a protease inhibitor often eventually experience treatment failure.

Since patients in the developed world are generally prescribed cocktails of therapeutic drugs, not all HIV-1 infections originate with a wild type but with drug sensitive strains, from which drug resistance inevitably emerges. As such, with the increase in prevalence of drug resistant strains, there comes an increase in infections that actually begin with drug resistant strains. Infections with pre-existing drug resistance immediately reduce the drug options for drug treatment and emphasize the importance of drug resistance information to optimize initial therapy for these patients.

Moreover, as the number of available antiretroviral agents has increased, so has the number of possible drug combinations and combination therapies. It is therefore very difficult, if not impossible, for the physician to establish the optimal combination for an individual. Although there are many drugs available for use in combination therapy, the choices can quickly be exhausted and the patient can rapidly experience clinical progression or deterioration if the wrong treatment decisions are made. The key to tailored, individualized therapy lies in the effective profiling of the individual patient's virus population in terms of sensitivity or resistance to the available drugs. This requires the advent of truly individualized therapy.

There are certain solutions to this problem currently in use.

Phenotyping directly measures the actual sensitivity of a patient's pathogen or malignant cell to particular therapeutic agents. However, this can be slow, labor-intensive and thus expensive.

A second approach to measuring resistance involves genotyping tests that detect specific genetic changes (mutations) in the viral genome which lead to amino acid changes in at least one of the viral proteins, known or suspected to be associated with resistance. Although genotyping tests can be performed more rapidly, a problem with genotyping is that there are now over 100 individual mutations with evidence of an effect on susceptibility to HIV-1 drugs and new ones are constantly being discovered, in parallel with the development of new drugs and treatment strategies. The relationship between these point mutations, deletions and insertions and the actual susceptibility of the virus to drug therapy is extremely complex and interactive. An example of this complexity is the M184V mutation that confers resistance to 3TC but reverses AZT resistance. The 333D/E mutation, however, reverses this effect and can lead to dual AZT/3TC resistance.

Sophisticated interpretation is therefore required to predict what the net effect of these mutations might be on the susceptibility of the virus population to the various

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therapeutic agents. Custom algorithms such as rules-based computer algorithms have provided some assistance, for example, see International patent application WO01/79540. An overview of this type of technique is presented in Figure 1.

Beerenwinkel et al., PNAS (Jun 2002), 99(12), pp 8271-8276; Schmidt et al., AIDS (Aug 2000), 14(12), pp 1731-1738; and Sevin et al., Journal of Infectious Diseases, (Jul 2000), 182(1), pp 59-67; disclose methods for quantitating the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV based on different algorithms such as, respectively, decision trees, a rule-based approach and statistical analyses such as cluster analysis, recursive partitioning, linear discriminant analysis. In Schmidt et al., AIDS Reviews, 4(3), pp 148-156, further methods are reviewed.

Meisel et al., Therapeutic Drug Monitoring (Feb 2001), 23(1), pp 9-14; and Meisel et al., Pharmacogenetics (1997), 7(3), pp 241-246; disclose a method for predicting the metabolic activity phenotype from the mutation pattern of the NAT-2 gene by multiple linear regression analysis. The linear regression model describes a quantity R_S, the metabolic ratio built up by an error term (first term) and a sum of products built up from a mutation factor multiplied by a mutation-dependent resistance coefficient. However, given the nature of the NAT-2 genotypic patterns, the above methods do not consider the relationship between point mutations within a genotypic pattern. In particular, the quantitative prediction methods proposed are merely an addition of independent variables where effects such as antagonism or synergy between point mutations, insertions or deletions are not taken into account.

There remains a continuing need for the quantitative prediction of HIV drug susceptibility from viral genotype. In particular, there is a need for quantitative prediction methodologies like linear regression modelling which can grasp the complexity of the HIV-1 genotypic-to-phenotypic dynamics, i.e. combinatorial effects such as antagonism and synergism. Furthermore, because the majority of HIV patients have now been exposed to drug cocktails, it is thought that the disease-causing retroviruses tend to spontaneously generate mutations that have often co-evolved. This makes the analysis of which mutations are responsible for resistance to which drugs almost impossible using currently available techniques. It also means that mutations that contribute to resistance are being overlooked using the currently available analysis techniques.

It is therefore an aim of the present invention to provide methods for improving the interpretation of genotypic results.

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It is a further aim of the invention to provide methods for determining (or predicting) a phenotype based on a genotype.

It is also a further aim of the invention to provide methods for predicting the resistance of an HIV variant of a particular genotype to a therapy or a therapeutic agent.

5 It is also an aim of the invention to predict resistance of a patient to therapy.

It is also an aim of the invention to provide methods to assess the effectiveness or efficiency of a therapy or to optimize a patient's therapy.

It is also an aim of the invention to identify novel HIV-1 mutations that are associated with resistance to particular drug therapies or combination therapies.

SUMMARY OF THE INVENTION

A solution to these problems involves new methods for measuring drug resistance by correlating genotypic information with phenotypic drug resistance profiles measured experimentally.

According to a first aspect of the invention, there is provided a method for quantitating the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV, said method comprising the steps of:

a) performing a linear regression analysis using data from a dataset of matching genotypes and phenotypes, whereby the log fold resistance, pFR, is modelled as the sum of all the individual resistance contributions for each of the mutations or combinations of mutations that occur in HIV according to the following equation;

$$pFR = \beta_A M_A + \beta_B M_B + \dots + \beta_Z M_Z + \varepsilon$$

wherein each individual resistance contribution is calculated by multiplying a mutation factor, M_A , M_B , ..., M_Z , for each mutation or combination of mutations by a resistance coefficient β_A , β_B , ..., β_Z ;

wherein the mutation factor assigned to each mutation or combination of mutations reflects the degree to which that mutation or combination of mutations is present in the HTV strain and, if present, to which degree the mutation is present in a mixture;

wherein each resistance coefficient reflects the contribution of the mutation or combination of mutations to the fold resistance exhibited by the strain;

and wherein the error term ϵ , represents the difference between the modelled prediction and the experimentally determined measurement.

This method involves a data driven technique for quantitative drug susceptibility prediction. This method uses a multiple linear regression model to estimate coefficient

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values that accurately reflect the contribution made by a particular HIV mutation or combination of mutations to resistance to a particular drug. Repeating the method for each candidate therapeutic drug allows the compilation of a global picture of drug resistance exhibited by a particular HIV strain.

This method has allowed the identification of mutations hitherto unrecognized as having an effect on drug resistance in HIV. The method also allows the identification of primary (single mutations) and secondary (the co-occurrence of two mutations) or higher order terms resistance-associated mutations for new and existing drugs, which embrace the antagonistic and synergistic phenomena. Accordingly, a further aspect of the invention provides a method of identifying a mutation that affects the degree of drug resistance exhibited by an HIV strain using a method according to the first aspect of the invention.

The method of the first aspect of the invention is also advantageous over current methods since it allows the quantitative, purely data-driven, objective_assessment of the contribution of mutations and combinations of mutations to drug resistance. The method also allows the de-convolution of the individual contribution made by particular mutations to the drug resistance phenotype. Unlike existing methods, the method is able to correct for correlating mutations that on the face of it appear to affect drug resistance, but which in fact only correlate in their occurrence with resistance causing mutations and are themselves phenotypically silent.

The method has allowed the design of an automated computational technique for the prediction of the drug resistance profile possessed by a particular HIV strain infecting a patient. The methods thus allow the determination of a patient phenotype without having to perform any phenotypic testing whatsoever. This has clear ramifications for the bespoke design, optimisation and assessment of strategies for individual patient therapy based upon the genotype of the infecting agent.

The invention also provides diagnostic kits for performing each of the methods of the invention described herein.

30 **DESCRIPTION**

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In any population of HIV variants, there is a wide distribution of drug resistance phenotypes for any particular drug, ranging from hyper-susceptibility to strong resistance (see Figure 2). The expression "drug resistance phenotype" means the resistance of an HIV virus to a tested therapy, therapeutic agent or drug. The term "resistance" as used herein, pertains to the capacity of resistance, sensitivity, susceptibility, hyper-susceptibility or effectiveness of a therapy against a disease. The term "therapy" includes but is not limited to a drug, pharmaceutical, or any other

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compound or combination of compounds that can be used in therapy or therapeutic treatment of HIV. This distribution of drug resistance reflects the large number of different genotypes that are present in the population. Some variants may only have one mutation that is correlated with drug resistance, whilst others will have several or numerous such mutations, each of which may impart its own contribution to the drug resistance phenotype.

Adding an additional level of complication are the phenomena of antagonism, synergy and enhancement, where certain mutations may add to or detract from the effect of other mutations in a manner not predictable from studying the effects of the individual mutations alone. Highly correlated mutations are also problematic. These are mutations that almost always co-occur in a strain, but only one of the mutations actually has an effect on drug resistance. For example, when one of these 2 mutations has an effect on resistance and the other mutation does not (this mutation might for example be highly correlated with the resistance mutation because it affects the replication rate of the virus), the effect can erroneously be assigned to either one of the mutations.

Examples of mutations known or suspected to influence the sensitivity of HIV to drug therapy may be found on the internet at http://hiv-web.lanl.gov; http://hivdb.stanford.edu/hiv/; or http://www.viral-resistance.com.

In HIV, two sections of the genome are generally studied: Protease (PR) and Reverse Transcriptase (RT). The methods of the present invention can equally be applied to other sections of the HIV genome such as Integrase (IN). A mutation is presented as a number referring to the position in the protein, followed by the amino acid(s) on that position, if it differs from the amino acid in the HXB2 HIV reference. In the terms included above, the mutations are represented as "A", "B", ..."Z".

Mixtures reflect the diversity of the HIV population in a sample. It means that on that position two subsets of the population have a different amino acid. Mixtures are denoted by separating amino acids with the '/' character: 65K/R (mixture of 'K' and 'R' at position 65).

When more than two amino acids are found on a certain position in subsets of the population, the dummy amino acid 'X' is used.

Insertions are denoted by adding the insert position behind a dot: 69.2S (an insert of 'S' at insert position 2). Deletions are denoted by a minus sign: 69-.

Examples of mutations present in the RT domain of HIV conferring resistance to a reverse transcriptase inhibitor include 69C, 69V, 69T, 75A, 1011, 103T, 103N, 184T,

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188H, 190E, 219N, 219Q, 221Y, 2211, and 233V. Additional examples of mutations present in the protease (PR) domain of HIV conferring resistance to a reverse transcriptase inhibitor include 24M, 48A, and 53L. A mutation may affect resistance alone or in combination with other mutations.

For the purposes of the invention, the mutations identified should be associated with resistance or susceptibility to drug therapy, for example an antiretroviral drug. The degree to which a particular mutation pattern may affect resistance may be determined by one of skill in the art, for example, using the phenotypic resistance monitoring assay such as, the ANTIVIROGRAM® (Virco, Belgium) (see WO97/27480). In this methodology, resistance is determined with respect to a laboratory reference strain HIVLAI/IIIB. The difference in IC₅₀ (the concentration of drug required to reduce the virus' growth in cell culture by 50%) between the patient sample and the reference viral strain is determined as a quotient. This fold change in IC₅₀ is reported and indicative of the resistance profile of a certain drug. Based on the changes in IC₅₀, cut-off values have been established to distinguish a sample from being sensitive or resistant to a certain drug.

Various projects are underway to compile data relating to the correspondence of certain mutations with drug resistance phenotype, and these generally lead to the generation of relational databases of tables that illustrate the matching genotype / resistance phenotype for various antiretroviral drugs. Such databases bring together the knowledge of both a genotypic and phenotypic database. The phenotypic database contains phenotypic resistance values for HIV to at least one therapy, preferably multiple drug therapies. For example, the phenotypic resistance values of tested HIV viruses, with a fold resistance determination compared to the reference HIV virus (wild type).

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The dataset used herein is a dataset developed by the Applicant, which consists of a set of matching genotype / phenotype measurements with possible multiple phenotype measurements per genotype. However, any similar dataset may be used, provided that there are sufficient entries for each genotype / phenotype measurement for the data to be significant. In the Virco dataset, the mutations are defined relative to HXB2 at amino acid level.

The phenotypes are presented as pFR values, where pFR is equal to -log(FR), where FR denotes the Fold Resistance. Negative pFR values thus denote resistance and positive values denote hyper-susceptibility. For example, a pFR value of -1.0 is equal to 10-fold resistance. An example of the pFR distribution for Saquinavir (SQV) is shown in Figure 3. Figure 4 shows the pFR distribution for the "48V" mutation on

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SQV. It is clear from this that the 48V subset does not behave the same as the whole dataset.

The problems of unwanted correlations between mutations where not all correlated mutations contribute to the drug resistance phenotype are illustrated in Figure 6. Here, the left hand panel shows the pFR distribution for the 71I mutation. When the effects of mutations 48V and 84V are removed (right hand panel), the pFR in the distribution of variants is markedly increased (less drug resistance).

According to the invention, the predicted fold resistance of an HIV strain of a particular genotype may be calculated by summing the individual resistance contributions for each of the mutations or combinations of mutations in the mutation pattern of that genotype. The method uses linear regression models, so that the phenotype prediction, pFR is calculated in the following equation (1):

$$pFR = \beta_A M_A + \beta_B M_B + \dots + \beta_Z M_Z + \varepsilon$$

The independent variables M_A , M_B , ..., M_Z , are referred to herein as mutation factors, each of which reflects the degree to which the mutation or combination of mutations is present in the HIV strain and, if present, whether or not the mutation is present in a mixture.

The resistance coefficients β_A , β_B , ..., β_Z represent the contribution to the total pFR prediction for each single mutation.

Each mutation factor M_i thus represents the presence or absence of the corresponding mutation and each coefficient β_i represents the contribution to the pFR change for that specific mutation.

The mutation factor may take into account 1st order terms (single mutations) as well as 2nd order terms (the co-occurrence of two mutations) and in general nth order terms. For example, 2nd order terms take the form:

$$\beta_{AB}^{M}_{AB}$$

The independent variable MAB represents the co-occurrence of mutations A and B and the coefficient βAB represents the synergy or antagonism between mutations A and B. When the mutation factor embraces n^{th} order terms, thus the co-occurrence of two ore more mutations, the terms take the form:

$$\beta_n M_n$$

wherein M_n represents the co-occurrence of one mutation with other one or more mutations—such as duplets, triplets, quadruples, etc. and the coefficient β_n represents

the synergy or antagonism between the one mutation with the other one or more mutations; thus n will apply for instance to the combinations of mutations, AB, ABC, ABCD, BC, ACD, etc.

As such, the linear regression model may take the following equation:

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$$pFR = \beta_A M_A + \beta_B M_B + \beta_n M_n + \dots + \beta_Z M_Z + \varepsilon$$

Higher order terms affect for interactions between mutations:

- antagonism or reversal: positive pFR shifts for mutation couples.
- synergy or enhancement: extra negative *pFR* shift for mutation couple.
- For example, consider the following (artificial) model:

Mutation	Coefficient
84V	-0.46
50V	-0.92
54M	-0.64
88S	0.63
90M	-0.16
46L	-0.19
46L & 84V	-0.09

Consider a virus with following mutations: 3I, 46L, 84V and 90M. Applying equation (1), this virus will have a pFR prediction:

$$pFR = \beta_{46L}.1 + \beta_{84V}.1 + \beta_{90M}.1 + \beta_{46L,84V}.1 = -0.9$$

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or almost 8-fold resistance, i.e. $pFR = -\log FR$.

Note that in the model F and A are synergistic since their co-occurrence decreases the pFR by an extra -0.09.

The error term is ε , which is the difference between the prediction and the measurement. This error term contains both the measurement error on the phenotype measurement and a model error (if the underlying model has higher order terms that are not taken into account in the regression model).

Mutation factors for single mutations (M_A , M_B , ..., M_Z) are calculated as follows:

if the mutation is present in the HIV strain, a positive mutation factor is assigned;

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if the single mutation is not present, the mutation factor assigned is zero; if the single mutation is present in a mixture, an averaged positive mutation factor is assigned.

Conveniently, mutation factors range between 0 and 1 where 0 means not present and 1 means present. Values between 0 and 1 means that the mutation is present in a mixture. Accordingly, a positive mutation factor is assigned the value 1.

Mixtures are modelled as causing the average shift of its constituent mutations. Since methods for the quantitation of the precise proportions of mixtures to wild type are expensive and time-consuming, mixture with wild type may conveniently be treated as causing half the pFR shift of the resistance mutation (mutation factor = 0.5). However, as the skilled reader will appreciate, a more precise mutation factor may be assigned if the true proportion in the mixture is known.

Mutation factors for double mutations (MAB etc.) are calculated as follows;

if both the mutations are present in the HIV strain, a positive mutation factor is assigned (conveniently, the value 1);

if neither of the mutations are present, the mutation factor assigned is zero;

if both mutations are present and one mutation is present in a mixture, an averaged positive mutation factor is assigned (conveniently, 0.5);

if both mutations are present in a mixture, a reduced averaged positive mutation factor is assigned (in this example, 0.25). The factor 0.25 is the product of the M-factors of both the single constituent mutations. This is the result of the assumption that these mixtures are independent of each other. Of course, this is an approximation, since in a real blood sample, the mixtures are not independent of each other. For example, if only 2 viruses were present, virus A (no mutations) for 70% and virus B (mutations 46I and 84V) for 30%, then a mixture would be detected on both positions 46 and 84. If these concentrations were known, it would be possible to fine tune the mutation factor of 0.25. If this information is not available, the best statistical guess is 0.5*0.5: this being the average value that would be measured for the mutation couple being present for a population of samples that have these mixtures on 46 and 84 in all possible concentrations.

Similarly, the mutation factors for triple mutations ($MABC\ etc.$) shall be calculated as follows:

if the three mutations are present in the HIV strain, a positive mutation factor is assigned (conveniently, the value 1);

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if neither of the three mutations are present, the mutation factor assigned is zero; if the three mutations are present and one mutation is present in a mixture, an averaged positive mutation factor is assigned (conveniently, 0.5);

if the three mutations are present and two mutations are present in a mixture, a reduced averaged positive mutation factor is assigned (in this example, 0.25). The factor 0.25 is the product of the M-factors of the two single constituent mutations present in the mixture;

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if the three mutations are present in a mixture, a reduced averaged positive mutation factor is assigned (in this example, 0.125). The factor 0.125 is the product of the M-factors of the three single constituent mutations.

The calculation of the mutation factors for higher order terms shall take the same principle.

Calculation of the resistance coefficient (β_A , β_B , ..., β_Z , β_AB) is performed by evaluating the dataset for the drug phenotype reported for each mutation or combination of mutations.

The problem of unwanted correlations has been discussed above. Unwanted correlations are removed according to the methods of the invention.

One way to do this is to use an algorithm that has been developed by the inventors to track the change in pFR as the effects of individual mutations or combinations of mutations are removed from the dataset. The effect of each mutation or combination of mutations is thus separated out. The methodology follows mutation trajectories towards the global average as the effects of individual mutations or combinations of mutations are removed. The steps are as follows:

- a) calculate average pFR for all mutations with a sufficient count in the database to be significant;
- b) determine the extremes (maximum, minimum), and select the mutation with the pFR furthest away from the global average;
- c) remove all virus strains that have the selected mutation and reiterate from step a);
- d) stop when the selected mutation in step b) has an average pFR that approximates to the global average.

In this manner, mutations that do not cause resistance, but which are often present with mutations that do cause resistance will have a higher average pFR (less resistance). Removing the virus strains with a certain resistance causing mutation results in an increase of the average pFR for correlating mutations.

A suitable threshold at which a count in the database becomes sufficiently significant will be apparent to the skilled reader and will be dependent on the database size. For example, thresholds of 5, 10, 15, 20, 25, 30 or more may be suitable. In the examples discussed herein, a threshold of 20 times was used.

- By an "average pFR that approximates to the global average" is meant that the average pFR is within a fraction of the standard deviation of the remaining population. A convenient fraction ranges between about 0.3 and 0.5.
 - A comparison of the change in the global average pFR with the change in the average pFR for selected mutations with increasing iterations of the algorithm is shown in
- Figure 7. Figure 8 shows an example, where the average pFR for 71I (unwanted correlation) jumps up as a result of removing from the dataset virus strains that have "71I & 84V" and "48V" mutations.

An alternative, analogous methodology for removing unwanted correlations is as follows; this is an extension of the mutation trajectories algorithm discussed above.

- 15 The steps of this method are as follows:
 - a) calculate correlation coefficient between all mutations (with a sufficient count in the database) and the pFR;
 - b) determine the extremes (maximum, minimum), and select the mutation with the highest (absolute value of) correlation coefficient;
- c) calculate a linear model for the pFR with the selected mutation(s) (from step b), all previous iterations);
 - d) take the residue (pFR minus the predicted value from the model);
 - e) calculate correlation coefficient between all mutations (with a sufficient count in the database) and the residue;
- 25 f) determine the extremes (maximum, minimum), and select the mutation with the highest (absolute value of) correlation coefficient;
 - g) calculate a linear model for the pFR with the selected mutation(s) (from step f), all previous iterations); and
 - h) reiterate from step d);
- i) stop when the selected mutation in step g) has a correlation coefficient that approximates to zero.

As with the mutation trajectories algorithm described above, the effect of mutations that do not themselves cause resistance, but which are often present with mutations that do cause resistance, is excluded and thus does not distort the real values.

In a more preferred methodology for removing unwanted correlations, a stepwise selection regression may be applied, which method selects the variable with the highest effect. The steps of this method are as follows:

- a) perform a first order regression from the list of mutations that occur in the dataset;
- b) calculate the p-value for all mutations;
- c) select the mutation with the lowest p-value and add it to the model;
- 10 d) re-calculate the regression model;
 - e) reiterate from step b);

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f) stop when the re-calculation of the p-values of step b) gives no significant values anymore.

Usually, this methodology is run in statistical software packages, which iteratively model the residue from the previous regression as the dependent variable.

$$pFR = Intercept_{1} + \beta_{A}M_{A} + \varepsilon_{1}$$

$$\varepsilon_{1} = Intercept_{2} + \beta_{B}M_{B} + \varepsilon_{2}$$

$$\varepsilon_{2} = Intercept_{3} + \beta_{C}M_{C} + \varepsilon_{3}$$

The p-value for a given mutation is the probability of rejecting the true null hypothesis, where the null hypothesis is defined as follows: the coefficient of that parameter equals zero. In other words, the p-value is the probability that the real coefficient for a certain parameter is zero, while the model predicts a coefficient different from zero.

The expression "significant value" for the p-values refers to the mutations with a p-value that is lower than the threshold selected by the user. This threshold may be determined as follows: the user shall create linear models for a whole range of combinations of p-values (a p-value for the first-order iteration and a p-value for the

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second-order iteration). For each combination, the mean squared error (MSE) of the corresponding model is calculated on unseen data. The combination of p-values that results in the model with the lowest MSE, i.e. the combination for which the model gives the best predictions, is chosen.

In further preferred embodiments of the invention, problems of small datasets for particular mutations or combinations of mutations are dealt with by applying the method recursively to the set of virus strains that exhibit those particular mutations or combinations of mutations.

In still further preferred embodiments of the invention, the following additional correlations are taken into account:

- multiple entries of the same virus strain (or virus strains grown from the same stock solution) that cause unwanted correlations;
- censored values in genotype / phenotype database (for example, EC₅₀ value = '< 1μM'). These are phenotypes beyond the assay range, thus when the phenotypic value is smaller than the measurable range, a '<'-censor is applied to that value. Analogously, a '>'-censor is applied to the value, if it is higher than the measurable range.

Preferably, censored values are dealt with by attempting to construct a model that is consistent from extrapolations. Censored values are thus modeled by replacing the censored value by a maximum likelihood estimation, assuming knowledge of the standard deviation of the measurement error.

A preferred technique for the generation of a maximum likelihood estimation is as follows:

- a) calculate a linear regression model without censored values;
- 25 b) use the phenotypic measured value V_0 as if the censor was "=", e.g. when a result is expressed as $-\log FR < 4$, we will treat V_0 as $-\log FR = 4$;
 - c) look at the prediction P from the model and apply either:

Case '<'-censor:

- $P < V_0 0.798 \sigma$ (center of gravity of half Gaussian distribution)
 - o Remove value from training data for next iteration
- V_0 − 0.798 $\sigma \le P < V_0$
 - o Use $V' = V_0 0.798$ σ for next iteration
- $V_0 \le P$

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o Use V' centre of gravity of tail (<V) of a normal distribution N (P, σ) as value for next iteration.

Case '>'-censor:

- $P > V_0 + 0.798$ σ (center of gravity of half Gaussian distribution)
- o Remove value from training data for next iteration
 - V_0 + 0.798 σ ≥ P > V_0
 - o Use V' = $V_0 0.798 \sigma$ for next iteration
 - $V_0 \ge P$

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- O Use V' centre of gravity of tail (>V) of a normal distribution N (P, σ) as value for next iteration.
- d) calculate a linear regression model and for the censored values in the linear regression model, either remove the data-point from the training set, or use V' instead of the censored phenotypes measurement, as described in step c);
- e) re-iterate from step b) until the prediction converges.
- Accordingly, for each iteration, when the prediction and measurement contradict, censored values are taken into account. When the prediction and measurement are strongly consistent, censored values are disregarded, on the basis that no further information is provided and their inclusion has no additional value.
 - In one preferred embodiment of this aspect of the invention, the number of calculations necessary in the linear regression analysis may be reduced. The computational power and memory requirement that is currently generally available is insufficient to allow a full second order model to be evaluated for a large dataset, based on all possible single mutations and second order terms, since the number of terms increases quadratically with the number of mutations considered. This number increases with a larger dataset since more rare mutations are in a large database.
 - In order to reduce the amount of terms, a first order regression may be performed from the list of mutations that occur in the dataset above a threshold number of times. A suitable threshold at which a count in the database becomes sufficiently significant will be apparent to the skilled reader and will be dependent on the database size. For example, thresholds of 5, 10, 15, 20, 25, 30 or more may be suitable. In the examples discussed herein, a threshold of 20 times was used. The significant terms from this first order regression are withheld and the list of these terms is then used to perform a second order regression. In the second order regression only the single mutations and combinations of mutations are used that were found significant in the first order model.

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Again, a threshold significance will be apparent to the skilled reader – an example is if the probability that the real value of the term is 0, is smaller than 0.001.

For example, a first order regression performed on the matching genotype / phenotype dataset for indinavir (34,445 measurements) for those mutations that occur at least 20 times results in a first order model that withholds a list of 94 single mutations that are considered significant.

This list is then used as a starting list for a second order regression. It should be noted that it may be advantageous to exclude certain very common mutations from the calculation. 3I is one example. The reason is that a mutation must occur at least a threshold number of times and the inverse also has to be true: the count of viruses not having the mutation 3I or the couple not 3I and another mutation should also be above the threshold value (e.g. 20). Taking this into account results in excluding 3I from the regression in practice.

In the second order regression, all the single mutations and all couples of mutations from the list are used as potential terms. The significant terms are then withheld by the regression algorithm.

According to a further aspect of the invention, there is provided a method of calculating the quantitative contribution of a mutation pattern to the drug resistance phenotype exhibited by an HIV strain, said method comprising the steps of:

- a) obtaining a genetic sequence of said HIV strain;
 - b) identifying the pattern of mutations in said genetic sequence, wherein said mutations are associated with resistance or susceptibility to drug therapy; and
- c) calculating the fold resistance of the HIV strain as compared to the wild type HIV strain by performing a linear regression analysis, whereby the log fold resistance, pFR, is modelled as the sum of all the individual resistance contributions for each of the mutations or combinations of mutations that occur in said HIV strain according to the following equation;

$$pFR = \beta_A M_A + \beta_B M_B + \dots + \beta_Z M_Z + \varepsilon$$

wherein each individual resistance contribution is calculated by multiplying a mutation factor, MA, MB, ..., MZ, for each mutation or combination of mutations by a resistance coefficient βA , βB , ..., βZ ;

wherein the mutation factor assigned reflects the degree to which the mutation or combination of mutations is present in the HIV strain and, if present, to which degree the mutation is present in a mixture;

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wherein each resistance coefficient reflects the contribution of the mutation or combination of mutations to the fold resistance exhibited by the strain;

and wherein the error term ϵ , represents the difference between the modelled prediction and the experimentally determined measurement.

As the skilled reader will appreciate, the fold resistance of the HIV strain may be calculated using any one of the embodiments of the invention referred to above.

In the first step of this method, the genetic sequence of an HIV strain should be obtained. Normally, this will be the genetic sequence of an HIV strain with which a patient is infected, although the sequence may be a theoretical sequence, for example for purposes of *in silico* modelling.

The method may thus be used as a diagnostic method for predicting the fold resistance exhibited by a particular HIV strain with which a patient is infected. According to other preferred embodiments, the method may be used for assessing the efficiency of a patient's therapy or for evaluating or optimising a therapy. The method may be performed for each drug or combination of drugs currently being administered to the patient so as to obtain a series of drug resistance phenotypes and thus to assess the effect of a plurality of drugs or drug combinations on the predicted fold resistance exhibited by the HIV strain with which the patient is infected.

A "patient" may be any organism, particularly a human or other mammal, suffering from HIV or AIDS or in need or desire of treatment for such disease. A patient includes any mammal and particularly humans of any age or state of development.

To obtain an HIV strain from a patient, a biological sample will need to be obtained from the patient. A "biological sample" may be any material obtained in a direct or indirect way from a patient containing HIV virus. A biological sample may be obtained from, for example, saliva, semen, breast milk, blood, plasma, faeces, urine, tissue samples, mucous samples, cells in cell culture, cells which may be further cultured, etc. Biological samples also include biopsy samples.

The genetic sequence of an HIV strain may be evaluated by a number of suitable means, as will be clear to those of skill in the art. Most suitable will be techniques that allow for specific nucleic acid amplification, such as the polymerase chain reaction (PCR), although other techniques such as restriction fragment length polymorphism (RFLP) analysis will be equally applicable.

Nucleic acid sequencing then allows the analysis of the mutation pattern in a particular nucleic acid sequence, either by classical nucleic sequencing protocols e. g. extension chain termination protocols (Sanger technique; see Sanger F., Nicher., Coulson Λ .

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Proc. Nat. Acad. Sci. 1977, 74, 5463-5467) or chain cleavage protocols. Such methods may employ such enzymes as the Klenow fragment of DNA polymerase I, Sequenase (US Biochemical Corp, Cleveland, OH), Taq polymerase (Perkin Elmer), thermostable T7 polymerase (Amersham, Chicago, IL), or combinations of polymerases and proofreading exonucleases such as those found in the ELONGASE Amplification System marketed by Gibco/BRL (Gaithersburg, MD). Preferably, the sequencing process may be automated using machines such as the Hamilton Micro Lab 2200 (Hamilton, Reno, NV), the Peltier Thermal Cycler (PTC200; MJ Research, Watertown, MA) and the ABI Catalyst and 373 and 377 DNA Sequencers (Perkin Elmer). Particular sequencing methodologies have been developed further by companies such as Visible Genetics. Any of the novel approaches developed for unraveling the sequence of a target nucleic acid, either now or in the future will be perfectly applicable to the analysis of sequence in the present invention (including but not limited to mass spectrometry, MALDI-TOF (matrix assisted laser desorption ionization time of flight spectroscopy) (see Graber J, Smith C., Cantor C. Genet. Anal. 1999, 14, 215-219) chip analysis (hybridization based techniques) (Fodor S P; Rava R P; Huang X C; Pease A C; Holmes C P; Adams C L Nature 1993, 364, 555-6) It should be appreciated that nucleic acid sequencing covers both DNA and RNA sequencing.

Once the genetic sequence of the HIV strain is known, the pattern of mutation must be identified in the sequence. The term "mutation" as this is used herein, encompasses both genetic and epigenetic mutations of the genetic sequence of wild type HIV. A genetic mutation includes, but is not limited to, (i) base substitutions: single nucleotide polymorphisms, transitions, transversions, substitutions and (ii) frame shift mutations: insertions, repeats and deletions. Epigenetic mutations include, but are not limited to, alterations of nucleic acids, e. g., methylation of nucleic acids. One example includes (changes in) methylation of cytosine residues in the whole or only part of the genetic sequence. In the present invention, mutations will generally be considered at the level of the amino acid sequence, and comprise, but are not limited to, substitutions, deletions or insertions of amino acids.

The "control sequence" or "wild type" is the reference sequence from which the existence of mutations is based. A control sequence for HIV is HXB2. This viral genome comprises 9718 bp and has an accession number in Genbank at NCBI M38432 or K03455 (gi number: 327742).

Identifying a mutation pattern in a genetic sequence under test thus relates to the identification of mutations in the genetic sequence as compared to a wild type sequence, which lead to a change in nucleic acids or amino acids or which lead to altered expression of the genetic sequence or altered expression of the protein encoded

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by the genetic sequence or altered expression of the protein under control of said genetic sequence.

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A "mutation pattern" comprises at least one mutation influencing sensitivity of HIV to a therapy. As such, a mutation pattern may consist of only one single mutation. Alternatively, a mutation pattern may consist of at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine or at least ten or more mutations. A mutation pattern is thus a list or combination of mutations or a list of combinations of mutations. A mutation pattern of any particular genetic sequence may be constructed, for example, by comparing the tested genetic sequence against a wild type or control sequence. The existence of a mutation or the existence of one of a group of mutations can then be noted.

One way in which this may be done is by aligning the genetic sequence under test to a wild type sequence noting any differences in the alignment. Typical alignment methods include Smith-Waterman (Smith and Waterman, (1981) J Mol Biol, 147: 195-197), Blast (Altschul et al. (1990) J Mol Biol., 215(3): 403-10), FASTA (Pearson & Lipman, (1988) Proc Natl Acad Sci USA; 85(8): 2444-8) and, more recently, PSI-BLAST (Altschul et al. (1997) Nucleic Acids Res., 25(17): 3389-402). It may in some circumstances be preferable to generate alignments using a multiple alignment program, such as ClustalW (Thompson et al., 1994, NAR, 22(22), 4673-4680). Other suitable methods will be clear to those of skill in the art (see also "Bioinformatics: A practical guide to the analysis of genes and proteins" Eds. Baxevanis and Ouellette, 1998, John Wiley and Sons, New York). A practical example of multiple sequence alignment is the construction of a phylogenetic tree. A phylogenetic tree visualizes the relationship between different sequences and can be used to predict future events and retrospectively to devise a common origin. This type of analysis can be used to predict a similar drug sensitivity for a sample but also can be used to unravel the origin of different patient sample (i. c. the origin of the viral strain).

In this manner, therefore, the pattern of mutations in the genetic sequence can be identified, wherein said mutations are associated with resistance or susceptibility to drug therapy exhibited by the HIV strain tested. The mutation pattern may influence sensitivity to a specific therapy, e. g., a drug, or a group of therapies. The mutation pattern may, for example, increase and/or decrease resistance of the HIV strain to a therapy. Particular mutations in the mutation pattern, may also, for example, enhance and/or decrease the influence of other mutations present in the genetic sequence that effect sensitivity of the HIV strain to a therapy.

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The invention further relates to a diagnostic system as herein described for use in any of the above described methods. An example of such a diagnostic system, for quantitating the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by an HIV strain, comprises:

5 a) means for obtaining a genetic sequence of said HIV strain;

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- b) means for identifying the mutation pattern in said genetic sequence as compared to wild type HIV;
- c) means for predicting the fold resistance exhibited by the HIV strain using any one of the methods described above.
- The means for predicting the fold resistance are preferably computer means.

A still further aspect of the invention relates to a computer apparatus or computer-based system adapted to perform any one of the methods of the invention described above, for example, to quantify the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV, or to calculate the quantitative contribution of a mutation pattern to the drug resistance phenotype exhibited by an HIV strain.

In a preferred embodiment of the invention, said computer apparatus may comprise a processor means incorporating a memory means adapted for storing data; means for inputting data relating to the mutation pattern exhibited by a particular HIV strain; and computer software means stored in said computer memory that is adapted to perform a method according to any one of the embodiments of the invention described above and output a predicted quantified drug resistance phenotype exhibited by an HIV strain possessing said mutation pattern.

A computer system of this aspect of the invention may comprise a central processing unit; an input device for inputting requests; an output device; a memory; and at least one bus connecting the central processing unit, the memory, the input device and the output device. The memory should store a module that is configured so that upon receiving a request to quantify the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV, or to calculate the quantitative contribution of a mutation pattern to the drug resistance phenotype exhibited by an HIV strain, it performs the steps listed in any one of the methods of the invention described above.

In the apparatus and systems of these embodiments of the invention, data may be input by downloading the sequence data from a local site such as a memory or disk drive, or alternatively from a remote site accessed over a network such as the internet. The sequences may be input by keyboard, if required.

The generated results may be output in any convenient format, for example, to a printer, a word processing program, a graphics viewing program or to a screen display device. Other convenient formats will be apparent to the skilled reader.

The means adapted to quantify the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV, or to calculate the quantitative contribution of a mutation pattern to the drug resistance phenotype exhibited by an HIV strain will preferably comprise computer software means. As the skilled reader will appreciate, once the novel and inventive teaching of the invention is appreciated, any number of different computer software means may be designed to implement this teaching.

According to a still further aspect of the invention, there is provided a computer program product for use in conjunction with a computer, said computer program comprising a computer readable storage medium and a computer program mechanism embedded therein, the computer program mechanism comprising a module that is configured so that upon receiving a request to quantify the individual contribution of a mutation or combination of mutations to the drug resistance phenotype exhibited by HIV, or to calculate the quantitative contribution of a mutation pattern to the drug resistance phenotype exhibited by an HIV strain, it performs the steps listed in any one of the methods of the invention described above.

The invention further relates to systems, computer program products, business methods, server side and client side systems and methods for generating, providing, and transmitting the results of the above methods.

The invention will now be described by way of example with particular reference to a specific algorithm that implements the process of the invention. As the skilled reader will appreciate, variations from this specific illustrated embodiment are of course possible without departing from the scope of the invention.

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BRIEF DESCRIPTION OF THE FIGURES

- Figure 1: Overview of measured /predicted phenotype handling
- Figure 2: Phenotype distribution of ritonavir for matching G/P samples in Virco database
- 5 Figure 3: pFR distribution for saquinavir
 - Figure 4: Distribution of pFR for '48V' mutation on saquinavir
 - Figure 5: Distribution of pFR for '48V' mutation on saquinavir (expanded)
 - Figure 6: Removing unwanted correlations
 - Figure 7: global mutation trajectories
- Figure 8: mutation trajectories for 71I
 - Figure 9: Example of genotypes, mutations relative to HBX2
 - Figure 10: Example of phenotype analysis for ritonavir
 - Figure 11: Higher order interaction between mutations 82A and 84V
 - Figure 12: Illustration of iterative procedure for censored values
- Figure 13: Linear regression model identifies mutations included in IAS list. Mutations marked with an * are also identified by a regression on a 5% subset of the data
 - Figure 14: Linear regression model identifies additional mutations previously described in the literature
 - Figure 15: Predicted versus measured log(FC)
- Figure 16: Comparison between linear regression model and decision trees.
 - Figure 17: Histogram of population left after removing all virus strains during the iterations
 - Figure 18: Trajectory of mutation 18H
 - Figure 19: Residues as a function of the measured values
- 25 Figure 20: Histogram of the residues as a function of the measured values
 - Figure 21: Histogram of the residues as a function of the measured values after 6 parameters were taken into account

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EXAMPLES

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Example 1: Methodology

1.1 Introduction

This exercise involved the generation of a list of key mutations for each of the following drugs: Indinavir, Ritonavir, Saquinavir, Nelfinavir, Amprenavir, Lopinavir, Zidovudine, Didanosine, Zalcitabine, Stavudine, Abacavir, Lamivudine, Tenofovir, Nevirapine, Delavirdine and Efavirenz.

The obtained list of key mutations is derived from a linear regression model using single mutations and couples of mutations as independent variables. The dataset used for this analysis is an export of the Virco dataset at 2003/02/01 from the *vircomining* tables. Table 1 shows the matching geno/pheno counts for each drug (*each* phenotype measurement for a genotype counts as one measurement).

Table 1: matching geno/pheno counts

Drug	Count	Drug	Count	Drug	Count
Amprenavir	29,508	Lamivudine	34,395	Delavirdine	32,450
Indinavir	34,445	Abacavir	32,744	Efavirenz	32,601
Lopinavir	7,410	Stavudine	34,420	Nevirapine	34,738
Nelfinavir	34,470	Zalcitabine	34,539		31,750
Ritonavir	34,502	Didanosine	34,227		
Saquinavir	34,543	Tenofovir	14,591		
		Zidovudine	33,575		

15 **1.2 Dataset**

The used dataset consists of a set of matching genotype/phenotype measurements with possible multiple phenotype measurements per genotype. The mutations are defined relative to HXB2 at amino acid level. The phenotypes are presented as pFR values, which is equal to $-\log (FR)$, where FR denotes the Fold Resistance.

Negative pFR values denote resistance and positive values denote hyper-susceptibility. For example, a pFR value of -1.0 is equal to 10-fold resistance.

1.3 Linear regression

For example, consider the following (artificial) model:

Table 2

Mutation	Coefficient
84V	-0.46
50V	-0.92
54M	-0.64
88S	0.63
90M	-0.16
46L	-0.19
46L & 84V	-0.09

Consider a virus with following mutations: 3I, 46L, 84V and 90M. This virus will have a *pFR* prediction:

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$$pFR = \beta_{46L}.1 + \beta_{84V}.1 + \beta_{90M}.1 + \beta_{46L,84V}.1 = -0.9$$

or almost 8-fold resistance. Note that in the model 46L and 84V are synergistic since their co-occurrence decreases the pFR by an extra -0.09. Note that in the model the mutation 3I shows no resistance coefficient assigned and therefore it is not considered in the pFR prediction equation.

Figure 9 shows an example of four different genotypes (mutations relative to HBX2), whilst Figure 10 shows an example of phenotype analysis for RTV performed according to the method of the invention.

15 1.4 Model creation

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Using our facilities, it was computationally infeasible to calculate a full second order model on all possible mutations and second order terms, since the number of terms increases quadratically with the number of mutations considered.

E.g. for APV:

- Total number of occurring mutations and couples of mutations: 19,074
- mutations and couples with each at least 20 measurements: 4,107

In order to reduce the amount of terms, a first order regression was performed from the list of mutations that occur at least 20 times in the dataset. The significant terms from this regression were withheld and the list of these terms -except mutation 3I for some of the PI's, was used to perform a second order regression. A term is called significant if the probability that the real value of the term is 0, is smaller than 0.001. A mutation

must occur at least 20 times and the inverse also has to be true: the count of viruses not having the mutation 3I or the couple not 3I and another mutation should also be at least 20. Taking this into account results in excluding 3I from the regression in practice. In the second order regression only the single mutations and couples of mutations are used that were significant in the first order model.

1.5 Example of model creation: Indinavir

A first order regression is performed on the matching geno/pheno dataset (34,445 measurements of which 28,480 unique Virco IDs) for those mutations that occur at least 20 times. The resulting first order model withholds a list of 94 single mutations that are considered significant. This list (except 3I) is used as a starting list for a second order regression. In this second order regression, all the single mutations and all couples of mutations from the list are used as potential terms. The significant terms are withheld by the regression algorithm.

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1.6 The impact of cross-drug correlation on the significance level of mutations
Correlation between mutations that cause resistance to different drugs, has an impact on
the confidence of the coefficient for this mutation. One of the effects is that for nonnucleoside reverse transcriptase inhibitors (NNRTIs) and nucleoside reverse
transcriptase inhibitors (NRTIs), some non-relevant mutations for that drug appear as

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significant (though with a coefficient close to 0), because drug resistance to the drug is correlated with drug resistance to drugs that bind at a different place.

Note that this is only a problem for *interpretation* of the model. For *prediction* of the Fold Resistance, the resulting model remains a good *pFR* predictor.

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1.7 Effects of second order terms

Antagonism

Table 3

Parameter	pFR shift	Count
•••		1
82A & 84V	0.43	395
•••	•••	1
82A	-0.27	4845
•••	•••	
84V	-0.26	3531
•••		1

Second order terms can indicate a *synergy* or an *antagonism*. In the example above, the occurrence of either 82A or 84V cause a resistance shift, but the co-occurrence of both mutations almost completely cancels out the effect of both mutations and shifts to susceptible ranges. In case both mutations are present, the net *pFR* shift is only -0.43, while it is -0.26 or -0.27 if only one of the mutations are present. This is an example of strong antagonism.

Synergy

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Table 4

Parameter	pFR shift	Count
•••		
24I	-0.22	1022
•••		
24I & 73S	-0.48	30
•••	•••	·
73S	-0.20	2216
•••	•••	

In this example, 24I and 73S both cause a resistance shift, but their co-occurrence causes a strong extra shift towards resistance. When only one of the mutations is present, the *pFR* shift is -0.20 or -0.22, but the presence of both mutations causes a *pFR* shift of -0.48. 24I and 73S are thus strongly synergistic in this example.

• Enhancement

Table 5

Parameter	pFR shift	Count
•••		•••
32I	0	821
•••	•••	•••
32I & 82A	-0.26	516
•••		
82A	-0.27	4845
•••	•••	

32I by itself does not contribute to resistance, but it increases the resistance for an 82A mutation. 32I enhances the effect of the 82A mutation.

An example of the effects of higher order interactions is shown in Figure 11.

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1.8 Highly correlated mutations

Highly correlated mutations (i.e. mutations that almost always co-occur in a strain) can affect the results of a regression analysis For example, when one of these 2 mutations has an effect on resistance and the other mutation does not (this mutation might for example be highly correlated with the resistance mutation because it affects the replication rate of the virus), the effect can be assigned to either one of the mutations. Unless this is compensated for, the regression model will assign the effect to that mutation that reduces the prediction error the most, which might not always be the mutation that is biologically responsible for the effect. Due to the correlation, it would otherwise not be possible to distinguish between these mutations.

Another effect that occurs due to correlation is when a mutation is highly correlated with a pair of mutations in which the first mutation is present.

Table 6

Parameter	pFR shift	Count
•••	•••	
58N	-1.47	108
•••		
58N & 77L	1.16	106
•••	•••	
77L	0	471
•••		

In the above example, 108 samples have a 58N mutation and out of these, 106 samples also have a 77L mutation. The effect of a pure 58N mutation can only be derived from the samples that have 58N and do not have 77L, which leads to higher uncertainty on the estimated pFR shift of the 58N mutation. The couple-term '58N & 77L' will compensate for a too low estimation of 58N by having a too high estimation for its pFR shift.

Techniques are provided in the description to deal with these effects. The algorithm developed by the inventors tracks the change in pFR as the effects of individual mutations or combinations of mutations are removed from the dataset.

A comparison of the change in the global average pFR with the change in the average pFR for selected mutations with increasing iterations of the algorithm is shown in Figure 7. Figure 8 shows an example, where the average pFR for 71I (unwanted correlation) jumps up as a result of removing from the dataset virus strains that have "71I & 84V" and "48V" mutations.

Example 2: Illustration of a stepwise regression with amprenavir

In a dataset of 31,292 matching genotypes and phenotypes for amprenavir; a stepwise first order regression was performed, and the first 11 iterations are shown here. The inventors selected as p-values:

5 p-value entry = 0.001

p-value stay ≈ 0.5

After the first iteration all the variables with mutations found more than 20 times were selected. In the next step the p-value was calculated, and then the algorithm picked the variable with the lowest p-value. In this case it was P084V, and then a model was built with this variable (84V). The model is shown in

Table 7.

Variable P084_V Entered: R-Square = 0.2477 and C(p) = 43445.43

Table 7

		Analysis of Varia	nce		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	1	2042.55324	2042.55324	10301.6	<.0001
Error	31289	6203.83650	0.19828		4.000
Corrected Total	31290	8246.38974			

Variable Parameter Estimate Standard Error Type II SS F Value Pr > FIntercept -0.00318 0.00269 0.27810 1.40 0.2363 P084 V -0.81754 0.00805 2042.55324 10301.6 <.0001

In the next iteration, we repeated the previous process except that the influence of 84V is now removed. In this second run, the mutation with the lowest p-values was P082A.

20 Table 8 below.

Variable P082_A Entered: R-Square = 0.3538 and C(p) = 32908.89

Table 8

		Analysis of Varia	nce		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	2	2917.39937	1458.69968	8564.44	<.0001

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		Analysis of Varia	nce		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Error	31288	5328.99038	0.17032		
Corrected Total	31290	8246.38974			

Intercept 0.07033 0.00269 116.28598 682.75 <.00	Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr>F
P082_A -0.47905 0.00668 874.84612 5136.47 <.00	Intercept	0.07033	0.00269			<.0001
P084 V -0.82827 0.00747 2007 (77)	P082_A	-0.47905	0.00668	874.84612		<.0001
	P084_V	-0.82827	0.00747	2095.67091	12304.3	<.0001

In the next iteration, we repeated the previous process except that the influence of 84V and 82A is now removed. In this third run, the mutation with the lowest p-values was P090M.

It may happen that after some iterations, a mutation which was found first significant is not significant anymore, and as such it is removed from the model.

The resistance coefficient is adjusted after every iteration because of the addition of a new variable to the regression. E.g. for P84V, the resistance coefficient β changes from -0.81754 to -0.82827.

The following iterations were done in the same way. The results obtained are here below enclosed.

Variable P090_M Entered: R-Square = 0.4245 and C(p) = 25885.06

Table 9

Analysis of Variance						
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F	
Model	3	3500.64130	1166.88043	7692.82	<.0001	
Error	31287	4745.74844	0.15168			
Corrected Total	31290	8246.38974				

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Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr > F

Parameter Estimate	Standard Error	Type II SS	F Value	Pr > F
0.13680	0.00276	373.40964		<.0001
-0.40431	0.00642	601.20353		<.0001
-0.64830	0.00762			<.0001
-0.33549	0.00541			<.0001
	0.13680 -0.40431 -0.64830	0.13680 0.00276 -0.40431 0.00642 -0.64830 0.00762	0.13680 0.00276 373.40964 -0.40431 0.00642 601.20353 -0.64830 0.00762 1097.72370	0.13680 0.00276 373.40964 2461.75 -0.40431 0.00642 601.20353 3963.52 -0.64830 0.00762 1097.72370 7236.89

Variable P033_F Entered: R-Square = 0.4729 and C(p) = 21082.65 Table 10

		Analysis of Varia	nce		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	4	3899.47454	974.86863	7016.41	<.0001
Error	31286	4346.91521	0.13894		
Corrected Total	31290	8246.38974			

Parameter Estimate	Standard Error	Time II SS	777.	
	Julian Biloi	Type II SS	F Value	Pr > F
0.14140	0.00264	398.56350	2868.58	<.0001
-0.60263	0.01125	398.83323	2870.52	<.0001
-0.35755	0.00621	460.87758	3317.07	<.0001
-0.60772	0.00733	954.28785		<.0001
-0.30702	0.00521			<.0001
	-0.35755 -0.60772	0.14140 0.00264 -0.60263 0.01125 -0.35755 0.00621 -0.60772 0.00733	0.14140 0.00264 398.56350 -0.60263 0.01125 398.83323 -0.35755 0.00621 460.87758 -0.60772 0.00733 954.28785	0.14140 0.00264 398.56350 2868.58 -0.60263 0.01125 398.83323 2870.52 -0.35755 0.00621 460.87758 3317.07 -0.60772 0.00733 954.28785 6868.28

Variable P046_I Entered: R-Square = 0.5110 and C(p) = 17296.94

Table 11

Analysis of Variance							
Source	DF	Sum of Squares	Mean Square	F Value	Pr>F		
Model	5	4213.90735	842.78147	6538.51	<.0001		
Error	31285	4032.48240	0.12890				
Corrected Total	31290	8246.38974					
							

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vanable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr > F
Intercept	0.15801	0.00257	489.12701	3794.77	<.0001
P033_F	-0.59310	0.01084	386.19290	2996.18	<.0001
P046_I	-0.32298	0.00654	314.43281	2439.45	<.0001
FUOZ A	-0.32721	0.00601	381.95629	2963.31	<.0001
P084_V	-0.53666	0.00721	714.53640	5543.55	<.0001
P090_M	-0.25737	0.00511	326.53882	2533.37	<.0001

Variable P047_V Entered: R-Square = 0.5287 and C(p) = 15544.70

Table 12

Analysis of Variance							
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F		
Model	6	4359.53390	726.58898	5848.07	<.0001		
Елгог	31284	3886.85584	0.12424				
Corrected Total	31290	8246.38974					

Variable Parameter Estimate Standard Error Type II SS F Value Pr > FIntercept 0.15941 0.00252 497.68905 4005.73 <.0001 P033_F -0.55617 0.01069 336.13339 2705.42 <.0001 P046_I -0.27816 0.00655 223.90478 1802.13 <.0001 P047_V -0.63671 0.01860 145.62656 1172.10 <.0001 P082_A -0.32763 0.00590 382.93722 3082.13 <.0001 P084_V -0.54696 0.00708 740.88446 5963.13 <.0001 P090 M -0.25573 0.00502 322.35883 2594.56 <.0001

Variable P046_L Entered: R-Square = 0.5409 and C(p) = 14326.32

Table 13

Analysis of Variance							
Source	DF	Sum of Squares	Mean Square	F Value	Pr>F		
Model	7	4460.84208	637.26315	5266.21	<.0001		
Error	31283	3785.54767	0.12101				
Corrected Total	31290	8246.38974					

Variable	D		 	· · · · · · · · · · · · · · · · · · ·	7
variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr>F
Intercept	0.16438	0.00249	526.67853	4352.36	<.0001
P033_F	-0.54921	0.01056	327.61200	2707.32	<.0001
P046_I	-0.31345	0.00658	274.55948	2268.90	<.0001
P046_L	-0.27769	0.00960	101.30817	837.19	<.0001
P047_V	-0.63940	0.01835	146.85706	1213.60	<.0001
P082_A	-0.26575	0.00620	222.00011	1834.56	<.0001
P084_V	-0.52972	0.00702	689.88420	5701.06	<.0001
P090_M	-0.24267	0.00498	287.89278	2379.09	<.0001

Variable P050_V Entered: R-Square = 0.5534 and C(p) = 13094.82

Table 14

Analysis of Variance							
DF	Sum of Squares	Mean Square	F Value	Pr > F			
8	4563.24026	570.40503	4844.61	<.0001			
31282	3683.14948	0.11774					
31290	8246.38974						
	8 31282	DF Sum of Squares 8 4563.24026 31282 3683.14948	DF Sum of Squares Mean Square 8 4563.24026 570.40503 31282 3683.14948 0.11774	DF Sum of Squares Mean Square F Value 8 4563.24026 570.40503 4844.61 31282 3683.14948 0.11774			

Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr > F

Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr>F
Intercept	0.16629	0.00246	538.66106	4575.00	<.0001
P033_F	-0.51815	0.01046	288.64295	2451.52	<.0001
P046_I	-0.29956	0.00651	249.44262	2118.58	<.0001
P046_L	-0.27934	0.00947	102.51249	870.67	<.0001
P047_V	-0.64754	0.01811	150.58308	1278.94	<.0001
P050_V	-0.82797	0.02808	102.39819	869.70	<.0001
P082_A	-0.25944	0.00612	211.32860	1794.87	<.0001
P084_V	-0.53915	0.00693	713.14511	6056.94	<.0001
P090_M	-0.24415	0.00491	291.39382	2474.89	<.0001

Variable P054_M Entered: R-Square = 0.5643 and C(p) = 12005.72

Table 15

Analysis of Variance					
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	9	4653.81644	517.09072	4502.38	<.0001
Error	31281	3592.57330	0.11485		
Corrected Total	31290	8246.38974			

Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr > F
Intercept	0.16633	0.00243	538.92765	4692.51	<.0001
P033_F	-0.46793	0.01049	228.56387	1990.14	<.0001
P046_I	-0.30134	0.00643	252.40077	2197.69	<.0001
P046_L	-0.28508	0.00935	106.71653	929.19	<.0001
P047_V	-0.55613	0.01818	107.50855	936.09	<.0001
P050_V	-0.84757	0.02774	107.23423	933.70	<.0001

Variable	Parameter Estimate	Standard Error	Type II SS	F Value	D- > E
P054_M	0.55110		- JPC H 55	1 Value	Pr > F
F U 34_1VI	-0.57119	0.02034	90.57618	788.66	<.0001
P082_A	-0.26050	0.00605	213.05431	1855.09	<.0001
P084_V	-0.53337	0.00685	697.31870	6071.64	<.0001
P090_M	-0.23454	0.00486	267.57375	2329.80	<.0001

Variable P054_L Entered: R-Square = 0.5740 and C(p) = 11047.41

Table 16

		Analysis of Vari	ance		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	10	4733.53615	473.35362	4214.95	<.0001
Error	31280	3512.85359	0.11230		
Corrected Total	31290	8246.38974			

Parameter Estimate	Standard Error	Type II SS	F Value	Pr>F
0.16756	0.00240	546.67520	4867.84	<.0001
-0.40056	0.01068	158.09630	1407.76	<.0001
-0.30750	0.00636	262.46469	2337.10	<.0001
-0.28311	0.00925	105.23686	937.08	<.0001
-0.52353	0.01802	94.83277	844.43	<.0001
-0.87024	0.02744	112.94107		<.0001
-0.45611	0.01712	79.71971		<.0001
-0.61432	0.02018	104.09593		<.0001
-0.26465	0.00598			<.0001
-0.51838				
-0.22560				<.0001
	-0.40056 -0.30750 -0.28311 -0.52353 -0.87024 -0.45611 -0.61432 -0.26465	0.16756 0.00240 -0.40056 0.01068 -0.30750 0.00636 -0.28311 0.00925 -0.52353 0.01802 -0.87024 0.02744 -0.45611 0.01712 -0.61432 0.02018 -0.26465 0.00598 -0.51838 0.00679	0.16756 0.00240 546.67520 -0.40056 0.01068 158.09630 -0.30750 0.00636 262.46469 -0.28311 0.00925 105.23686 -0.52353 0.01802 94.83277 -0.87024 0.02744 112.94107 -0.45611 0.01712 79.71971 -0.61432 0.02018 104.09593 -0.26465 0.00598 219.74203 -0.51838 0.00679 654.14626	0.16756 0.00240 546.67520 4867.84 -0.40056 0.01068 158.09630 1407.76 -0.30750 0.00636 262.46469 2337.10 -0.28311 0.00925 105.23686 937.08 -0.52353 0.01802 94.83277 844.43 -0.87024 0.02744 112.94107 1005.68 -0.45611 0.01712 79.71971 709.86 -0.61432 0.02018 104.09593 926.92 -0.26465 0.00598 219.74203 1956.68 -0.51838 0.00679 654.14626 5824.81

Table 17

		Analysis of Vari	ance		
Source	DF	Sum of Squares	Mean Square	F Value	Pr > F
Model	11	4810.94944	437.35904	3982.07	<.0001
Error	31279	3435.44030	0.10983		
Corrected Total	31290	8246.38974			

Variable	Parameter Estimate	Standard Error	Type II SS	F Value	Pr>F
Intercept	0.16239	0.00238	510.05266	4643.93	<.0001
P033_F	-0.39933	0.01056	157.11889	1430.54	<.0001
P046_I	-0.32619	0.00633	291.69184	2655.80	<.0001
P046_L	-0.29089	0.00915	110.98928	1010.54	<.0001
P047_V	-0.50941	0.01782	89.70713	816.77	<.0001
P050_V	-0.86163	0.02714	110.70009	1007.90	<.0001
P054_L	-0.46062	0.01693	81.29492	740.17	<.0001
P054_M	-0.61674	0.01995	104.91704	955.25	<.0001
P082_A	-0.25645	0.00592	205.77424	1873.53	<.0001
P084_V	-0.51212	0.00672	637.66342	5805.80	<.0001
P088_S	0.57942	0.02182	77.41329	704.83	<.0001
P090_M	-0.22099	0.00477	236.06351	2149.31	<.0001

Example 3: Dealing with censored values

In this example, the method developed by the inventors to deal with censored values was applied for the drug amprenavir. Firstly, a linear regression model without censored values was calculated, see the values 'APV_pFR' for iteration 0 in the Tables 18-21 below. The phenotypic measured value is < -2.083062017 for Virus strain 1, for the first iteration, the value V₀ is equal to the measured phenotype value APV_pFR, but the censor is considered as '='. This was followed by iteration nr. 1. Once the values were obtained from iteration 1, the inventors compared the prediction value P, e.g. for Virus strain 1, -2.362076895 with the phenotypic measured value <-2.083062017. For

virus strains 1, 2 and 3 below, the measured values have a censor '<'. For virus strains 4, the measured value has a censor '>'. The scenarios established by the inventors, when the case was '<'-censor, and when the case was '>'-censor, were applied. New linear regression models were calculated and for the censored values in the linear regression model, either the data-points from the training set were removed, as for virus strains 1, 2 and for virus strain 4 (iteration 2 and 3 only), or those data-points were used instead of the censored phenotypes measurement, as illustrated for virus strain 3 and for virus strain 4 (only iteration 1). The procedure was reiterated until the prediction converged.

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Virus strain 1: CASE '<' and P < V - 0.798 σ

 $V - 0.798 \sigma = -2.282562017$

Table 18

ITER	APV_censor	APV_pFR	Prediction P	value
0	<	-2.083062017		-2.083062017
11	<	-2.083062017	-2.362076895	
2	<	-2.083062017	-2.537358188	
3		-2.083062017		

15 Virus strain 2: CASE '<' and $P < V - 0.798 \sigma$

 $V - 0.798 \sigma = -2.108607374$

Table 19

ITER	APV_censor	APV_pFR	Prediction P	value
0	<	-1.909107374		-1.909107374
1	<	-1.909107374	-2.608207782	
2	i i	-1.909107374		
3	i i	-1.909107374		

Virus strain 3: CASE '<' and V <= P

20 V - 0.798 σ = -2.156216334

Table 20

ITER	APV_censor	APV_pFR	Prediction P	value
0	<	-1.956716334		-1.956716334
1	<	-1.956716334		-2.016673816
_2				-2.021216328
3		-1.956716334		

Virus strain 4: CASE '>' and $V < P \le V + 0.798 \sigma$ V + 0.798 σ = 0.714928642

Table 21

ITER	APV_censor	APV_pFR	Prediction P	value
0	>	0.515428642		0.515428642
11	>	0.515428642	0.671585565	
2			0.759236384	
3	>	0.515428642	0.770724812	

5 Example 4: Mutations trajectories

A methodology for removing unwanted correlations involved an algorithm developed by the inventors in which the change in pFR was tracked as the effects of individual mutations or combinations of mutations were removed from the dataset. The effect of each mutation or combination of mutations was separated out. The methodology followed mutation trajectories towards the global average as the effects of individual mutations or combinations of mutations were removed.

In a first step, the average pFR was calculated for all mutations with a sufficient count in the database to be significant, i.e. >20. In the Table below the first and last 10 mutations of a list of 368 mutations is shown with the corresponding calculated average pFR.

Table 22

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		T
Label	Count	Average pFR
54M	239	-1.35421631044816
50V	129	-1.28868671545329
47V	281	-1.27082426054035
84A	21	-1.23536347179756
76V	187	-1.20328089469875
89V	269	-1.19273200093637
91S	36	-1.15103609689224
33M	22	-1.12268052165089
84C	21	-1.11529918764429
54L	327	-1.09959239280592
_		

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18L	65	0.12143300387582

1387	0.121561573218861
1261	0.134344491996282
406	0.140725379487564
86	0.154734960149477
20	0.159585638802379
22	0.184594303565984
21	0.202798973752203
51	0.213932584381636
204	0.494820688128991
	1261 406 86 20 22 21 51

The extremes were determined, and the mutation with the pFR furthest away from the global average was selected. In Table 22, the mutation selected was 54M.

Following, all virus strains that had the selected mutation were removed, in total it amounted to 283 virus strains: from 26738 to 26455. A new

Table 23 of average pFR was generated for the remaining mutations. The list below shows the first and last 10 mutations of the obtained results.

Table 23

New table of average per mutation:

Label	Count	Average pFR
50V	129	-1.28868671545329
84A	21	-1.23536347179756
76V	168	-1.13676536523909
47V	215	-1.12849822159876
91S	35	-1.12648669210251
33M	22	-1.12268052165089
84C	21	-1.11529918764429
54L	327	-1.09959239280592
89V	213	-1.05990043312845
33F	852	-0.988257531387902

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18L	65	0.12143300387582
69K	1384	0.125380073379929
89M	1256	0.141136938991525
33V	404	0.147380396015872

17E	86	0.154734960149477
63I	20	0.159585638802379
62M	22	0.184594303565984
198	21	0.202798973752203
15L	51	0.213932584381636
88S	202	0.494762597743788

The previous steps were reiterated. In Table 24 there is listed the mutations which were selected at different iteration counts.

Table 24

Count	Mutation selected
26738	54M
26455	50V
26296	84A
26272	918
26220	47V
25947	76V
25762	54L
25365	22V
25260	33F
24545	32I
24183	84V
21784	54S
21728	82F
21577	54T
21427	24F
21397	24I
20894	73T
20702	73C
20604	55R
20324	95F
20194	10R
20122	54A
20062	82C
20039	88S
19818	46L

Count	Mutation selected
19247	231
19190	58E
18909	67F
18877	73 S
18105	53L
17920	10F
17568	82A
16637	461
16215	54V
16028	35G
15996	90M

The procedure was stopped when the last selected mutation had an average pFR that approximated to the global average. In Figure 17, a histogram of the population left after removing all virus strains during the iterations is shown. In Figure 18, the trajectory of mutation 18H, which has no significant phenotype, demonstrates the underlying cause that a virus strain is resistant due to other mutations than 18H (54M, 76V, 33F, 84V).

Example 5: Highly correlated mutations

A methodology for removing unwanted correlations proceeded as follows. In a first stage, the correlation coefficient between all mutations (with a sufficient count in the database, i.e. >20) and the pFR was calculated. In Table 25 below there is listed the first and last 10 coefficients of a list of 202 coefficients that were calculated.

Table 25

NAME	correlation with APV pFR
P084_V	-0.51430323209603
P090 M	-0.510166810424169
P010 I	-0.43843660921923
P046_I	-0.430221405121849
P071_V	-0.400337497273138
P033_F	-0.385947043862637
P082_A	-0.343946685713202
P054_V	-0.339269018202483
P032_I	-0.275105400287351

NAME	correlation with APV pFR
P010_F	-0.263412991920486

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P065_D	0.044821230615045
P012_A	0.0476139542192145
P014_R	0.0485859961407927
P033_V	0.0571605219086003
P041_K	0.0713497764580897
P063_S	0.071416853283617
P030 N	0.080199177466301
P089_M	0.0904859904655573
P069_K	0.0926109439533826
P088_S	0.101920449277774

Consequently, the extremes were determined (maximum, minimum), and the mutation with the highest (absolute value of) correlation coefficient was selected. In this case was P084_V.

In the following step, a linear model for the pFR with the selected mutation(s) (from step 2, all previous iterations) was calculated. The predicted model obtained was Predicted pFR = $-0.844 * M_{84}$

Following, the residue was taken (pFR minus the predicted value from the model). In Figure 19 a graph of the residues as a function of the measured values is shown. In Figure 20, the same graph is represented in the form of histograms where the distribution of the residue may be observed.

Then, the correlation coefficient between all mutations with a sufficient count in the database and the residue was calculated. Results of the first and last 10 variable are shown here below. It will be observed that the order of the mutations had changed because the influence of mutations P084_V had been removed.

Table 26

NAME	Correlation with Residue
P082_A	-0.402088216120049
P090_M	-0.373018037350597
P033_F	-0.347337204147266
P032_I	-0.325689507827784

P046_I	-0.324078296352946
P010_I	-0.321343698855709
P054_V	-0.311246000758214
P071_V	-0.290388889036061
P047_V	-0.272571613897309
P046_L	-0.234148912637279

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P012_S	0.0358242687063381
P014_R	0.0401577992704191
P012_A	0.0442436406740132
P063_S	0.0484345347109348
P033_V	0.0537012592855092
P030_N	0.0548222080117112
P041_K	0.0659977808671604
P089_M	0.078693680117922
P069_K	0.0835173137603877
P088_S	0.110781153455097

Consequently, the extremes were determined again, and the mutation with the highest absolute value of the correlation coefficient was selected. The mutation selected now was P082_A.

A new linear model for the pFR with the selected mutation P082_A, was calculated. pFR = -0.782*M84 + -0.435*M82

After 6 iterations, the following resistance coefficients for a selected group of mutations was obtained.

Table 27

Parameter	β
P084_V	-0.512345233450494
P082_A	-0.247464800298171
P090_M	-0.156011849225004
P033_F	-0.532092050020052
P046_I	-0.205719880528639
P047_V	-0.656374746817602

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In Figure 21, a graph is represented in the form of histograms showing the residues after the 6 parameters were taken into account.

The reiterative procedure continued until the last selected mutation had a correlation coefficient that approximated to zero.

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Example 6: Results from linear regression modeling

In an initial test, genotypes and corresponding phenotypes determined for ritonavir (RTV) for 28,540 HIV-1 clinical isolates were used. The linear regression analysis identified 20/22 RTV resistance-associated mutations described in the IAS mutation list (all except 10F and 77I) (see Figure 13). Additional mutations whose effect on RTV susceptibility had been previously described (e.g. 73S/T/C, 84A/C and 88D) were also identified (Figure 14). Overall, 53 single mutations and 96 pairs of mutations were identified as having significant effect on susceptibility to RTV.

The predicted phenotype was compared to the measured phenotype in a leave-one-out cross-validation, demonstrating a root mean square error of 0.31 (logFR) (see Figure 15. The error rate of the linear modeling method [5.62% (sensitivity=93.0%, specificity = 95.4%)], compared favourably to a decision tree-based model [Beerenwinkel, PNAS 99, (2002) 8271-8276] [10.2% (sensitivity=89.8%, specificity=89.7%)] (see Figure 16).

The robustness of the algorithm as a function of the size of the input dataset was assessed using smaller subsets of data. Nine of 22 IAS resistance-associated mutations for RTV could be identified with subsets ≥ 5% (1600 isolates) of the original data. However, the accuracy of the predicted contribution of the mutations improved with increasing dataset sizes up to 50% of the original database (median standard error of the predicted contributions decreased 50%). Some secondary mutations (e.g. 10R, 32I, 82S) were identified as having a significant contribution to resistance only when the subset size reached a similar 50% level.

Example 7: Comparison of genotype-to-phenotype prediction for different Artificial Intelligence techniques

Analyses were performed on matching genotype/phenotype datasets for all 16 HIV inhibitors currently available. The matching genotype/phenotype datasets consisted of approximately 30,000 data points for most drugs except Lopinavir and Tenofovir.

As an example, the following results for Ritonavir were obtained:

A log(FR) root mean squared error of 0.31 (MSE=0.096).

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- A classification error of 5.6% (with regard to a standard cutoff applied by the inventors of FR=3.5).
 - (sensitivity=93.0%, specificity=95.4%).
- Regression model identified 53 single mutations and 96 pairs of mutations. 20 out of 22 mutations of the IAS list are confirmed in this model.

Table 28

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	r ²	CV-MSE * (in log(FR))	Classification error	Nr of samples	Coverage	Model inspection
Neural network (Lopinavir) ^{1,2}	0.88	n/a	n/a	1,322	100%	possible N
Support Vector Machine (Ritonavir) ⁴	0.79	0.176	n/a	652	100%	N
Support Vector Machines (Tibotec data, Ritonavir)	0.81	0.144	n/a	17,453	100%	N
Decision tree (Ritonavir) ³		Classification only	10.2%	469	100%	Y
Clustering (<i>Indinavir</i>) ⁵	n/a	n/a	15.3%	1,152	100%	N
Self Organizing Map (<i>Saquinavir</i>) ²		Classification only	15%	811 (38 matching)	84%	N
Linear regression modeling (<i>Virco</i> data, <i>Ritonavir</i>)	0.88	0.096	5.6%	34,502	100%	Y

- (*) CV-MSE: cross-validation mean squared error. Depending on the analysis, it is a leave-one-out or a 10-fold cross-validation
- [1] A 28-Mutation Neural Network Model that Accurately Predicts Phenotypic Resistance to Lopinavir (LPV)
- D Wang, R Harrigan and BA Larder, Antiviral Therapy 2001 (Supplement 1): 105
- [2] Predicting HIV Drug Resistance With Neural Networks
 S Drăghici, R Potter, Bioinformatics, Vol. 19 no. 1, 2003 (p. 98-107).
- [3] Geno2pheno: Interpreting Genotypic HIV Drug Resistance Tests
 N Beerenwinkel et al., Intelligent Systems in Biology, Nov/Dec 2001.
- [4] Geno2pheno: estimating phenotypic drug resistance from HIV-1 genotypes N Beerenwinkel et al., Nucleic Acids Research, 2003, Vol. 31, No 13.
- 20 [5] Predicting phenotype from genotype: a comparison of statistical methods
 A Foulkes et al.

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Neural networks, support vector machine and clustering techniques are no suitable candidates for a quantitative prediction system with the requirement that such a system should have a descriptive power in order to be inspectable by experts and to be able to motivate certain predictions to customers. Decision trees allow experts to have insight in how the decision process works. But decision trees require massive amounts of data to build complex trees, since the information in the data is not optimally used.

Based on the performed analyses and the linear regression modelling feasibility study, linear regression modelling seems to meet the requirements for a quantitative prediction system. Linear regression seems to outperform other examined techniques and offers the possibility of inspection to HIV-experts. Considering the lower error rates, the linear regression modelling allows the optimisation of drug therapy in patients.

Example 8: Comparison of the regression linear model with a rules-based algorithm

The regression linear model developed by the inventors was used to test the drug-15 associated resistance of one HIV-1 sample on nevirapine, delavirdine, and efavirenz. In parallel, the same sample was run in a rules-based algorithm methodology as described in WO01/79540.

The results were expressed in FR, or fold change in IC50 or EC50, relative to reference wild-type virus, which is the drug concentration at which 50% of the enzyme activity is inhibited, and is expressed in μM units. The FR cutoffs values for normal susceptible ranges were 8, 10, and 6, for nevirapine, delavirdine, and efavirenz, respectively.

The phenotypic antiviral experiment was taken as the gold-standard by the inventors. Said phenotypic antiviral experiment was perfonned as described in WO97/27480.

Results of the three methodologies are enclosed in Table 3 below. 25

> According to the results obtained by the rules-based algorithm, the patient's sample would be susceptible to all three drugs. However, when the results obtained by the linear regression model were considered, the sample showed resistance against nevirapine and efavirenz and susceptibility against delavirdine. These last results were confirmed by the phenotypic antiviral experiments.

Table 29

drug	Rules-based alg	Rules-based algorithm		Phenotypic antiviral experiment	
	Matches in database	FR	FR	FR	

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Nevirapine	321	3.6	176.1	>89
delavirdine	301	1.9	3.5	3.7
efavirenz	300	1.9	12.9	17.5